

MEDICARE FORM

Ilumya[™] (tildrakizumab-asmn) Injectable Medication Precertification Request

Page 1 of 2

(All fields must be completed and legible for precertification review.)

For Illinois MMP: FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business: Please use other form.

Note: Ilumya is non-preferred. Preferred products may vary based on indication. See section G below.

Please indicate:	Start of treatment: S Continuation of ther			1 1		on indication	n. See section G below
Precertification Re	equested By:			Phone:		Fax:	
A. PATIENT INFOR	MATION						
First Name:			L	_ast Name:			
Address:			C	City:		State:	ZIP:
Home Phone:		Work Phor	ne:		Cell Phone:		
DOB:	Allergies:				E-mail:		
Current Weight:	lbs_or	kgs	Height:	inches or	cm	S	
B. INSURANCE INF	ORMATION						
Aetna Member ID #:			Does patient have other coverage?				
Group #:			_ If yes, provide ID#: Carrier Name:				
Insured:		Insure	ed:				
C. PRESCRIBER IN	IFORMATION						
First Name:		Last	Name:		(Check On] D.O. 🗌 N.P. 🗌 P.A.
Address:	1			City:		State:	ZIP:
Phone:	Fax:	St Lic	; #:	NPI #:	DEA #:		UPIN:
Office Contact Nam	e:				Phone:		
Address: City: Phone:	d Physician's con Center Phone: ne:	ZIP:		Name:	Office harmacy	Retail Pha Other Other	ZIP:
Request is for: Ilur	mya (tildrakizumab-asm	n): Dose:		Frequency:			Code:
F. DIAGNOSIS INFO	DRMATION – Please indica	ate primary ICD Co	de and specify a	ny other where applica	ble.		
Primary ICD Code:		Secondary IC			Other ICD	Code:	
G. CLINICAL INFOR For Initiation Reque Note: Ilumya is nor Preferred products Yes No Has Yes No Has Yes No Has	RMATION – Required clinic ests (clinical documentation- preferred. Avsola and Re may vary based on indic the patient had prior therap the patient had a trial, intol the patient had a trial, intol Enbrel (etanercept) Repared Repared Repared Repared For the patient had a trial, intol Repared Repared Repared Repared Repared Repared Repared Repared Re	al information must on required for all emicade are prefe ation. by with Ilumya (tildr lerance, or contrain lerance, or contrain umira (adalimumab	t be completed in requests): rred for MA pla rakizumab-asmn idication to Avso idication to any co) Otezla (ap	ns. Enbrel, Humira, C) within the last 365 da la (infliximab-axxq) or l of the following? (select premilast) □ Skyrizi (i	Dertification requ Dtezla, and Skyr ys? Remicade (inflix t all that apply): risankizumab-rz	ests. rizi are preferre mab)? aa)	ed for MAPD plans.
diagnosis (select all	re are any other medical re that apply): Enbrel (etanercept) □ H						d for the patient's



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION (continued) - R	Required clinical information must be con	npleted in its <u>entirety</u> for all pred	certification requests.						
Plaque Psoriasis:									
Please indicate the severity of the patient's disease: mild moderate severe									
☐ Yes ☐ No Is there evidence that the disease is active?									
Yes No Is there clinical documentation of chronic disease?									
Yes ☐ No Is the patient a candidate for systemic therapy or phototherapy? → Please select: ☐ phototherapy ☐ systemic therapy ☐ phototherapy and systemic therapy									
Please select: phototherapy systemic therapy phototherapy and systemic therapy Please provide the patient's Psoriasis Area and Severity Index (PASI) score:									
	, , , <u> </u>								
Please indicate the percentage of body surface area affected by plaque psoriasis:%									
Set the plaque psonasis involve sensitive areas? If yes, please select. In fands Theet Thee Thee Sensitive areas a									
\square Yes \square No Was the trial with systemic conventional DMARD(s) (e.g., methodexate, accreting of cyclosponne) menecuve?									
☐ Yes ☐ No Are systemic conventional DMARDs contraindicated?									
Please select: acetretin cyclosporine methotrexate mycophenolate None of the above									
Please indicate the length of the medication trial: Less than 1 month 1 month 2 months 3 months or greater									
☐ Yes ☐ No Was the trial with phototherapy ineffective?									
\Box Yes \Box No Was the trial with phototherapy not tolerated?									
☐ Yes ☐ No Is phototherapy contraindicated?									
Please check all that apply: Psoralens (methoxsalen, trioxsalen) with UVA light (PUVA)									
UVB with coal tar or dithranol									
	UVB (standard or narrow band)								
	Home UVB								
] None of the above								
Please indicate the length of trial: 🗌 Less than 1 month 🔲 1 month 🔲 2 months 🔲 3 months or greater									
For Continuation of Therapy (clinical documentation required for all requests):									
Please indicate the length of time on Ilumya (tild									
Yes No Is this continuation request a result of the patient receiving samples of Ilumya (tildrakizumab-asmn)?									
Yes No Will Ilumya (tildrakizumab-asmn) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)?									
☐ Yes ☐ No Is there clinical documentation supporting disease stability?									
Yes No Is there clinical documentation supporting disease improvement?									
Yes No Does the patient have any risk factors for TB?									
└────────────────────────────────────									
└────────────────────────────────────									
Please enter the results of the TB test: positive positive punknown									
Yes No Has the patient received llumya (tildrakizumab-asmn) within the past 6 months?									
Yes No Does the patient have a documented severe and/or potentially life-threatening adverse event that occurred during or following the previous infusion?									
\rightarrow Yes \square No Could the adverse reaction be managed through pre-medication in the home or office setting?									
Please indicate the severity of the disease at baseline (pretreatment with Ilumya (tildrakizumab-asmn)): mild moderate severe									
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Require	d):		Date: / /						
Any person who knowingly files a request for a	authorization of coverage of a medical pr	ocedure or service with the inte	nt to injure, defraud or deceive any						
insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent									
insurance act, which is a crime and subjects such person to criminal and civil penalties.									

The plan may request additional information or clarification, if needed, to evaluate requests.